Origins and Affinities of Modern Humans: A Comparison of Mitochondrial and Nuclear Genetic Data

L. B. Jorde, M. J. Bamshad, W. S. Watkins, R. Zenger, A. E. Fraley, P. A. Krakowiak, K. D. Carpenter, H. Soodyall, T. Jenkins, and A. R. Rogers

¹Department of Human Genetics, Eccles Institute of Human Genetics, University of Utah Health Sciences Center, and ²Department of Anthropology, University of Utah, Salt Lake City; ³Department of Human Genetics, School of Pathology, The South African Institute for Medical Research, University of the Witwatersrand, Johannesburg; and ⁴Department of Anthropology, Pennsylvania State University, University Park

Summary

To test hypotheses about the origin of modern humans, we analyzed mtDNA sequences, 30 nuclear restrictionsite polymorphisms (RSPs), and 30 tetranucleotide short tandem repeat (STR) polymorphisms in 243 Africans, Asians, and Europeans. An evolutionary tree based on mtDNA displays deep African branches, indicating greater genetic diversity for African populations. This finding, which is consistent with previous mtDNA analyses, has been interpreted as evidence for an African origin of modern humans. Both sets of nuclear polymorphisms, as well as a third set of trinucleotide polymorphisms, are highly consistent with one another but fail to show deep branches for African populations. These results, which represent the first direct comparison of mtDNA and nuclear genetic data in major continental populations, undermine the genetic evidence for an African origin of modern humans.

Introduction

Genetic data can reveal much about human evolution. Earlier genetic studies of human populations were based on blood group and protein polymorphisms (Nei and Roychoudhury 1982; Cavalli-Sforza et al. 1988), but these systems may not be selectively neutral and do not directly reflect variation at the DNA level. MtDNA has been used extensively in studies of human origins (Cann et al. 1987; Vigilant et al. 1991; Rogers and Harpending 1992; Sherry et al. 1994; Horai et al. 1995), and most of these analyses have shown excess genetic diversity in African populations. This result has been interpreted as support for an African origin for modern *Homo sapiens*, although some disagree with this interpretation (Spuhler

Received March 8, 1995; accepted for publication June 1, 1995. Address for correspondence and reprints: Dr. Lynn B. Jorde, Department of Human Genetics, Eccles Institute of Human Genetics, University of Utah Health Sciences Center, Salt Lake City, UT 84112. Email: lbj@thor.med.utah.edu

© 1995 by The American Society of Human Genetics. All rights reserved. 0002-9297/95/5703-0001\$02.00

1993; Templeton 1993). Drawbacks of mtDNA are that it provides a limited amount of information about genetic variation (Nei and Livshits 1989) and that it may not be selectively neutral (Excoffier 1990). More recently, nuclear DNA polymorphisms, including restriction-site polymorphisms (RSPs) (Bowcock et al. 1991; Kidd et al. 1991) and highly variable minisatellite and microsatellite polymorphisms (Deka et al. 1991, 1995; Edwards et al. 1992; Mountain and Cavalli-Sforza 1994; Bowcock et al. 1994; Di Rienzo et al. 1994), have been used to infer human evolutionary history.

While nuclear and mtDNA data reveal some evolutionary patterns in common (Bowcock et al. 1994), the degree of concordance between these two types of data has not been closely scrutinized. In particular, no study has compared both types of genetic variation in the same individuals across major continental populations. We present the results of a direct comparison of mtDNA and nuclear variation in a series of 243 Africans, Asians, and Europeans. Our analyses show that nuclear and mtDNA data present discordant pictures of human population affinities.

Material and Methods

The study population consists of 75 Africans (22 Sotho-Tswana, 14 Tsonga, 14 Nguni, 15 San, 5 Biaka Pygmies, and 5 Mbuti Pygmies), 78 Asians (12 Cambodians, 17 Chinese, 19 Japanese, 6 Malay, and 9 Vietnamese, 2 Koreans, and 13 individuals of mixed Asian ancestry), and 90 Europeans (20 unrelated French members of the CEPH kindreds and 70 unrelated Utah males whose ancestries are traced almost exclusively to Great Britain, Denmark, Norway, and Sweden [O'Brien et al. 1994]; the latter are consequently termed "northern Europeans"). The Sotho-Tswana, Tsonga, and Nguni are all Bantu speakers who are thought to have diverged from one another during the past 1,000–2,000 years. Informed consent was obtained from all subjects whose blood was drawn at the University of Utah.

African, Asian, and European DNAs were prepared from blood or transformed lymphoblast cell lines. DNA extractions were performed by established methods (Bell et al. 1981; Sykes 1983). Genotypes for 30 tetranucleo-

tide short tandem repeats (STRs) and 30 RSPs were analyzed using PCR. Genomic DNA sequences were amplified in 1× buffer (10 mM Tris, pH 8.3; 50 mM KCl; and 1.5 mM MgCl₂) using 20 ng of template genomic DNA, 50 µM dNTPs, 20 pmol of each primer, and 1 U Tag DNA polymerase in a total reaction volume of 25 μl. Samples were cycled 30 times in a Perkin-Elmer 9600 PCR machine. For all STR systems, one primer was endlabeled using γ^{32} P-ATP and polynucleotide kinase. One picomole of radiolabeled primer was added to the standard PCR reaction above. PCR products for the STR systems were separated by denaturing polyacrylamide gel electrophoresis and visualized by autoradiography. PCR-amplified segments for each of the RSPs were digested for 2 h by adding 10 U of the appropriate restriction enzyme in 10 μ l of 1× restriction enzyme buffer to each sample. Restriction fragments were separated by agarose gel electrophoresis (3% NuSieve) and visualized by ethidium bromide staining.

The polymorphisms of each major type (RSPs and STRs) are unlinked, and nearly all are located on separate chromosome arms. Genome Data Base (GDB) ID numbers for RSP markers are G00-178-350, G00-181-665, G00-206-704, G00-197-514, G00-180-400, G00-178-648, G00-251-590, G00-185-229, G00-171-808, G00-177-841, G00-177-108, G00-187-676, G00-285-016, G00-181-821, G00-180-411, G00-250-584, G00-178-649, G00-181-402, G00-182-559, G00-180-743, G00-196-856, G00-177-381, G00-210-937, G00-180-968, G00-182-529, G00-192-312, G00-196-905, G00-181-184, G00-182-305, G00-185-174. STR loci used in this analysis are D1S407, D1S399, D2S273, D3S1537, D3S1545, D4S1525, D4S1530, D5S580, D6S400, D6S393, D7S620, D7S623, D8S499, D8S384, D9S249, D9S762, D10S526, D10S516, D10S525, HRAS1 (GDB ID G00-187-026), VWFII (GDB ID G00-177-640), D14S119, D15S195, D16S485, D17S919, D18S390, D19S403, D19S400, D20S161, and D20S428.

The mtDNA region corresponding to hypervariable sequence-2 (HVS-2) in Vigilant et al. (1989) was PCR-amplified as described by Bamshad et al. (in press). After extensive sequencing of HVS-2 in 20 individuals, a 200-bp region corresponding to bases 71–270 of the Cambridge Reference Sequence (Anderson et al. 1981) was found to contain 90% of the polymorphisms in the region and was thus examined in all individuals. The mtDNA sequence and the allele frequencies for the STRs and RSPs are given in the appendix.

Data were analyzed at two levels of population subdivision: major continental populations (African, Asian, and European) and the 13 subpopulations listed within these major populations. Allele frequencies for each RSP and STR system were estimated directly by gene counting. Heterozygosity for each of these systems was estimated as $1 - \Sigma x_i^2$, where x_i is the estimated frequency of the *i*th allele in the system. Standard errors of these estimates were obtained by using equation 8.7 in Nei (1987). Nucleotide diversity for the mitochondrial sequence was measured as $(n/(n-1))\Sigma x_ix_j\pi_{ij}$, where n is the number of individuals, x_i is the frequency of the ith mtDNA haplotype in the population, and π_{ij} is the proportion of nucleotides that differ between the ith and jth mtDNA haplotypes (Nei 1987). The standard error of this estimate was estimated using equation 10.7 of Nei (1987). The proportion of genetic variance attributable to population subdivision was estimated using the G_{ST} statistic (Wright 1965; Nei 1987). Selective neutrality of the mtDNA data was evaluated using tests devised by Tajima (1989) and Rogers (in press).

For the nuclear polymorphisms, genetic distances between pairs of populations were estimated using Nei's standard distance (Nei 1987). This measure assumes an infinite-alleles model of mutation. Genetic distances for the STRs were also estimated using a new measure formulated by Shriver et al. (1995), which weights distances by the difference in the number of repeat units and thus assumes a stepwise mutation model. For the mtDNA sequence data, Kimura's (1980) two-parameter model was used to estimate nucleotide diversity between each pair of individuals. An empirically derived transition:transversion ratio of 30:1 was used (Bamshad et al., in press). The average diversity within and between populations was then estimated by equation 10.21 of Nei (1987).

The relationships among populations were depicted by neighbor-joining trees (Saitou and Nei 1987), using programs in the PHYLIP package (Felsenstein 1993). The neighbor-joining method tends to provide more accurate estimates of population affinities than some other methods because it does not assume equal evolutionary rates in each branch of the tree (Nei and Roychoudhury 1993). Two other tree-making methods (the Fitch-Margoliash least-squares method and the unweighted pair-group method using arithmetic averages) yielded similar results (data not shown). To assess the reliability of these trees, 1,000 bootstrap replicates were run for each set of systems.

The degree of congruence among the STR, RSP, and mtDNA distance matrices was evaluated using the Mantel matrix comparison test (Mantel 1967; Smouse et al. 1986). This test yields a product-moment correlation coefficient for each pair of distance matrices. An empirical significance level is obtained by repeatedly permuting the columns of one of the distance matrices and comparing the actual correlation coefficient with those generated from the permuted matrices. Each pair of matrices was permuted 10,000 times.

Results and Discussion

Genetic Diversity within Major Populations

The gene diversities for each major population are given in table 1. Europeans exhibit the highest level of

Table I

Gene Diversity (±Standard Error) for Each Major Population

	RSPs	STRs	mtDNA*
Africans	.322 ± .073	.769 ± .102	.031 ± .001
	.377 ± .052	.681 ± .074	.011 ± .001
Europeans Overall	$.432 \pm .032$	$.724 \pm .086$	$.010 \pm .001$
	$.377 \pm .018$	$.725 \pm .020$	$.020 \pm .001$

^a Diversity at the nucleotide level. The standard error estimates include sampling variance only. The standard errors for the mtDNA nucleotide diversity are larger if stochastic variance is also included.

heterozygosity for RSPs, while Africans have the lowest heterozygosity level. This pattern has been seen in other studies of RSPs (Bowcock et al. 1991) and could reflect the fact that most of these polymorphisms were first detected in European populations (Mountain and Cavalli-Sforza 1994). In contrast, the highest STR heterozygosity values are seen in Africans, and the lowest are seen in Asians. The average number of STR alleles per locus is highest in Africans (9.5), intermediate in Europeans (8.2), and lowest in Asians (8.1). These differences are not statistically significant using a one-way analysis of variance, however, and the standard errors of the heterozygosity estimates given in table 1 show that these values are not significantly different.

The gene diversity values based on mtDNA are similar in magnitude to values obtained in other studies (Horai et al. 1993). These diversity values differ significantly, with African diversity approximately three times higher than that of Asians and Europeans.

Application of Tajima's (1989) neutrality test to the mtDNA data shows that gene diversity in Europeans and Asians is considerably lower than predicted by neutral theory (D=-1.43 and -1.46, respectively). The African gene diversity is greater than predicted by neutral theory (D=0.60). However, these values lie within the 90% confidence limits of the neutral prediction, so the neutrality hypothesis is not rejected by this test. Rogers's test, which is based on the distribution of nucleotide differences within populations, rejects the hypothesis of mutation-drift equilibrium. This is consistent either with a rapid expansion of the human population or with a departure from selective neutrality.

Genetic Diversity between Populations

The genetic distances between each pair of continental populations are given in table 2. In each system—RSPs, STRs, and mtDNA—the Asian-European distance is smaller than the African-Asian and African-European distances. The Africans are more divergent for mtDNA sequence than for the two sets of nuclear polymorphisms: the ratio of distances involving Africans to those not involving Africans is approximately 2:1 for nuclear polymorphisms but 9:1 for the mtDNA sequence.

Table 2 also lists the $G_{\rm ST}$ values for each system. The $G_{\rm ST}$ estimate for RSPs, .107, indicates that ~11% of genetic variation can be attributed to subdivision at the level of major continental populations. This value lies within the 10%-15% range seen in other studies of RSPs (Bowcock et al. 1991), protein and blood group polymorphisms (Jorde 1980; Nei 1993; Cavalli-Sforza et al. 1994), and craniometric data (Relethford and Harpending 1994). The $G_{\rm ST}$ value obtained from STRs, .034, is substantially smaller than the value obtained from RSPs and likely reflects the relatively high STR mutation rate. It should be emphasized, however, that this estimate of $G_{\rm ST}$ does not take differences in STR allele sizes into account.

The G_{ST} estimate for mtDNA sequence, .199, is higher than the estimates based on nuclear polymorphisms. This difference may be attributed to the fact that the effective population size of mtDNA is one-fourth that of nuclear DNA (Birky et al. 1983), producing more rapid genetic drift for mtDNA polymorphisms. Previous G_{ST} estimates derived from mtDNA restriction site data range from .31 to .46 and are thus substantially higher than our estimate (Stoneking et al. 1990; Merriwether et al. 1991). In part, this may be because the control region, from which the present G_{ST} estimate was derived, has a much higher mutation rate than the remainder of the mitochondrial genome, from which the previous estimates were derived. In addition, the previous G_{ST} estimates were obtained using the method of Takahata and Palumbi (1985), which appears to yield inflated values (Harpending et al., in press).

Figure 1 displays a neighbor-joining tree based on mtDNA data in 13 subpopulations. All 13 subpopulations cluster into the appropriate major continental groups (Africans, Asians, and Europeans). Long branch lengths are seen for most of the African populations. This pattern has been observed in most other mtDNA analyses and has been a major component of the argument for an African origin of modern humans (Cann et al. 1987; Vigilant et al. 1991; Stoneking 1993). The non-African populations have comparatively short branch lengths, and the nodes separating these populations are very close to one another.

Figure 2, which presents a neighbor-joining tree estimated from RSP data, demonstrates pronounced clustering of each of the three major continental populations. As has been seen in other studies (Bowcock et al. 1994; Cavalli-Sforza et al. 1994), the Mbuti Pygmies have a long branch length; this may be caused by small effective population size. The Malay sample, which was derived from an aboriginal population, also has a long branch length. In contrast to the mtDNA tree, the African populations do not display markedly longer branch lengths in the RSP tree.

A neighbor-joining tree based on STRs is shown in figure 3. The distance measure of Shriver et al. (1995),

.010

.001

Genetic Distances between N	lajor Populations for Eac	h Major Genetic System	
	RSPs $(G_{ST} = .107)$	$STRs^a$ $(G_{ST} = .034)$	$mtDNA (G_{ST} = .199)$
African-Asian	154	.024	.009

Table 2

Genetic Distances between Major Populations for Each Major Genetic System

.124

.066

which assumes a stepwise mutation process, was used in estimating the tree. Figure 3 closely resembles figure 2, with populations clustering into major continental groups. This marked clustering argues against the hypothesis that the low $G_{\rm ST}$ values seen for STRs reflect interpopulation convergence resulting from forward-backward mutation (Bowcock et al. 1994). Instead, the high STR mutation rates produce low $G_{\rm ST}$ values because of a high level of within-group diversity relative to total diversity (Jin and Chakraborty, in press). Genetic

African-European

Asian-European

Chinese
French
63
687
N. European
26
Japanese

Malay

Cambodian
289
Vietnamese

Nguni
337
Sotho/Tswana
Biaka Pygmy

Tsonga

Mbutti Pygmy

Figure 1 Neighbor-joining tree based on mtDNA sequence data. In this and all subsequent figures, the numbers listed next to branch nodes indicate the no. of times that the bootstrapped replicates supported the branches to the right of the node. Absent numbers indicate that the configuration of the actual tree differed from that of the consensus tree at this node.

distances, however, can remain large and well defined, as observed in the present analysis.

.016

.010

An STR tree based on Nei's standard distance, which assumes an infinite alleles model, was less similar to the RSP tree and displayed less clear separation of continental population groups. This result provides indirect evidence that STR variation is modeled more accurately by a stepwise mutation process (such as replication slippage) than by an infinite alleles process (such as unequal chromosome exchange) (Shriver et al. 1993; Di Rienzo et al. 1994).

Table 3 contains the Mantel matrix correlations for each pair of distance matrices. These correlations give quantitative support to the visual interpretations of the

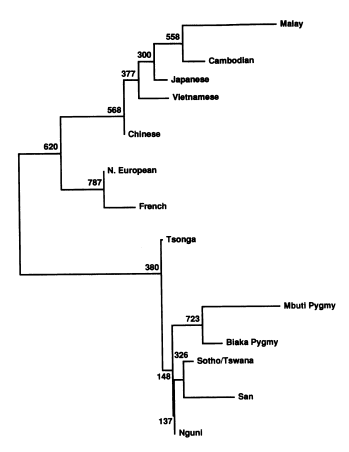


Figure 2 Neighbor-joining tree based on Nei's standard distance for 30 RSPs (Nei 1987). See legend to fig. 1.

^a The distances reported for STRs were estimated using the method of Shriver et al. (1995).

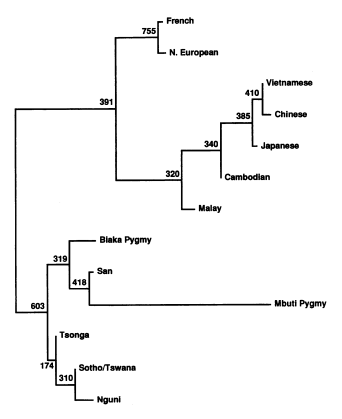


Figure 3 Neighbor-joining tree based on Shriver et al.'s (1995) distance for 30 STRs. Bootstrapping was done using each locus (rather than each allele) as the sampling unit. See legend to fig. 1.

neighbor-joining trees. In addition to the data sets described above, a distance matrix based on five trinucleotide repeat systems (Huntington disease, spinal bulbar muscular atrophy, spinal cerebellar ataxia type 1, myotonic dystrophy, and dentatorubral pallidoluysian atrophy [DRPLA]) is included. These data, which are described fully by Watkins et al. (in press), were generated on the same study population as the RSP, tetranucleotide, and mitochondrial systems. The tetranucleotide

and RSP distances yield a large and highly significant correlation ($P < 10^{-4}$), and the correlation is higher when the distance measure of Shriver et al. (1995) is used for the tetranucleotides. The trinucleotide repeat distances are also highly correlated with both of the other sets of nuclear systems ($P < 10^{-4}$). In contrast, all of the correlations involving mtDNA with nuclear systems are lower and either less significant or nonsignificant. The largest correlation is seen with RSPs (r = .512, P < .001), while the correlations with tetranucleotide systems are nonsignificant (P > .05) and the correlation with trinucleotide systems is low and marginally significant (r = .219; .04 < P < .05).

Implications for Modern Human Origins

Our mtDNA results are consistent with those of most other studies in showing increased gene diversity and long branch lengths for African populations. The nuclear DNA results are also consistent with previous studies of nuclear blood group and protein polymorphisms (Cavalli-Sforza et al. 1988; Nei and Roychoudhury 1993), RSPs (Bowcock et al. 1991), dinucleotide repeat polymorphisms (Bowcock et al. 1994; Di Rienzo et al. 1994; Deka et al. 1995), and a sample of four *Alu* polymorphisms (Batzer et al. 1994) in showing greater divergence for African populations than for other populations.

Although all data types suggest greater African divergence, the degree of divergence is less for the nuclear data than for the mtDNA data. In addition, while the nuclear RSP and STR results are highly concordant, there is less concordance between the mtDNA and nuclear results shown in figures 1–3. This has important implications for genetic inferences about modern human origins, since population history should affect mtDNA and nuclear DNA variation similarly. There are several possible reasons for this discordance:

I. Ascertainment bias in nuclear polymorphisms.—The mtDNA polymorphisms cannot be affected by an ascertainment bias, because they were obtained from DNA sequence ascertained uniformly in all populations. The

Table 3

Mantel Matrix Correlations for Each Pair of Genetic Distance Matrices

	RSP	STR (Nei's distance [Nei 1987])	STR (Shriver's distance [Shriver et al. 1995])	Trinucleotides
mtDNA	.512ª	.359	.274	.219 ^b
RSP		.655°	.769°	.747°
STR (Nei's distance)			.814°	.555°
STR (Shriver's distance)				.640°

NOTE.—Other correlations are nonsignificant.

 $^{^{}a}P < .001.$

 $^{^{\}text{b}}.04 < P < .05.$

 $^{^{\}circ} P < 10^{-4}$.

RSPs were detected primarily in Europeans, and the resulting bias may contribute to the excess European heterozygosity in the RSP data. This is unlikely, however, to be the sole cause of increased heterozygosity. A statistical analysis of ascertainment bias in two-allele systems demonstrates that this bias can account for no more than half of the excess European heterozygosity (A. R. Rogers and L. B. Jorde, unpublished information). A parallel analysis of systems with multiple alleles shows that ascertainment bias is negligible for the highly heterozygous STR polymorphisms. If ascertainment bias were responsible for the discordance observed here, one would predict that the RSPs, which presumably have the greatest degree of bias, would have the lowest correlation with mtDNA sequence. Instead, table 3 shows that they have the highest correlation. In addition, one of the expanded trinucleotide repeat polymorphisms, DRPLA, was ascertained in Asians. Yet this polymorphism yields a neighbor-joining tree with the same configuration as the tetranucleotide tree, with roughly equal branch lengths in Africans and Asians (Watkins et al., in press). Given these results, it is unlikely that ascertainment bias alone accounts for the discordance between mtDNA and nuclear DNA results.

- 2. Differences in substitution rates.—The substitution rate in the mtDNA HVS-2 region is estimated to be \sim 12.5% per nucleotide per million years, or 3×10^{-6} per nucleotide per generation (Sherry et al. 1994). This is considerably higher than that of the RSPs, whose substitution rate is estimated to be 10^{-7} – 10^{-9} per nucleotide per generation (Nei 1987; Bowcock and Cavalli-Sforza 1991). The mutation rates for tetranucleotide repeats are the highest of the three systems, estimated at 10^{-4} – 10^{-3} per generation (Edwards et al. 1992; Weber and Wong 1993). If substitution rates accounted for the discordance observed here, one would expect the two nuclear substitution rates to be most similar to one another. Instead, the mtDNA substitution rate is intermediate between those of the two sets of nuclear polymorphisms.
- 3. Inadequate sample sizes.—All population genetic analyses are necessarily based on a limited collection of individuals and polymorphisms. It is unlikely that the differences observed here are caused by a limited sample size, because all three nuclear data sets produce highly consistent results and the mtDNA results are consistent with those published previously using different population samples and different portions of the mtDNA genome (hypervariable sequence 1 and RSPs outside the D loop). Nonetheless, the degree of resolution of the mtDNA data is limited by the fact that the mtDNA molecule is effectively a single locus.
- 4. Differences in male-female migration rates or effective population sizes.—Because mtDNA is maternally inherited while nuclear DNA is biparentally inherited, differences in the two types of data could be produced by differences in male-female migration patterns or effective popula-

tion sizes. Higher mtDNA diversity between African populations would then require that male dispersal has been greater than female dispersal in African populations and/or that fewer females than males contributed to the African gene pool for a long period. Additional light will be shed on these possibilities as more information is gained about genetic variation in human Y chromosome DNA (Spurdle and Jenkins 1992).

5. Lack of selective neutrality in mtDNA.—An important assumption in inferring population history from genetic data is that genetic polymorphisms are selectively neutral. The RSPs and STRs used in this study occur primarily in noncoding DNA, and most appear to meet this assumption (Bowcock et al. 1991; Shriver et al. 1993). In contrast, mtDNA polymorphisms outside the noncoding D loop show a departure from neutrality (Excoffier 1990; Merriwether et al. 1991; Templeton 1993), with a relative lack of diversity seen in Asians and Europeans. The HVS-2 data analyzed here show a similar departure from neutrality in Asians and Europeans, although it is not statistically significant. These departures may reflect the action of natural selection, or they could be the result of past population expansions (Rogers and Harpending 1992; Rogers, in press). Since there is no recombination in the mitochondrial genome, natural selection on a coding gene will exert a substantial genetic "hitchhiking" effect, even on polymorphisms in the noncoding D loop. It is thus possible that the differences seen here in mtDNA and nuclear DNA may be produced by natural selection rather than population history.

Increased mtDNA diversity in Africans has been a linchpin of the argument that modern humans originated in Africa and then replaced existing archaic populations on other continents. Proponents of this view argue that since Africa is more diverse genetically, its population must be older (Stoneking 1993). However, diversity can be strongly affected by events in a population's history, such as the timing of major bottlenecks, and therefore does not necessarily reflect a population's age (Rogers and Jorde 1995). Our findings further compromise the diversity argument by showing that nuclear DNA trees lack the deep branches (and thus the excess genetic diversity) observed in mtDNA trees. These results do not disprove the African replacement hypothesis. However, they weaken the genetic evidence in its favor.

Acknowledgments

We wish to thank Li Jin for providing a computer program that calculates the Shriver et al. (1995) distance and Kenneth and Judith Kidd for providing Asian DNA samples. Elisabeth Manderscheid aided in blood sample collection. This research was supported by NSF grants BNS-8720330 and DBS-9310105, the Clinical Research Center of the University of Utah (NIH RR-00064), the Technology Access Center of the Utah Human Genome Project, and a supplemental equipment grant from the University of Utah.

Appendix

The data in figures A1-A3 are available in computerized format from the authors.

9	2000	000	0	0	0001	c	>	0	0	0009	0	3000	0	0001	3	> <	> 0	> (5	-	> 0	-	•	> •	2	1250	1250	1250	1250	1250	2500	0	1250	>	0	0	0	0	0	0	2500	0	2000	0	
표	3250	2750	250	0	750	c	•	0	0	833	833	Ξ	1667	2500		777	8	٠ ;	8/7			-																			1579				
	3471											280										5																			204				
	2778											1999										-													0	0	0	Ξ	0	0	0	0	0	929	,
	2002											900																													0				
	2763											6888										-																			263				
	1875											2940 3		•	•							-													0	0	0	0	0	0	294	0	0	0	
	1667											2083										> c													0	0	417	0	833	833	0	417	0	417	•
	2000											714 2	٠.									-																•			385				
	2500											000										-																			0				
	2500								• • •			1429																		• •											357				
TS	101	1429	2857	0	714	2500		0	2143	714	1071	714	2143	1786	1001	1701	ì	۰ د	>																0	0	0	357	0	357	714	714	357	0	
	2632							263	1316	1053	526	1842	2368	1053	200	3 5	000	8	>	263	3	-	•	ם כ	3 3	563	1842	2105	1842	1579	789	263	789	>	0	0	0	0	263	263	226	263	263	0	
Allele	S	9	7	∞	6	2	:	-	7	3	4	S	9	7	- o		٠ :	≘ :	=	-	٠,	۰ ۳	, •	4 v	٠,	9 1	7	œ	6	2	=	12	2 :	<u> </u>	_	7	3	4	2	9	7	∞	6	01	
Locus							D4S1525												0761630	D431330														1158580	Decer										
9	,	0	9	900	2000	4000	2000	0	0	0		0	۰ د	0	0	0	0	8750	1250	0	0	•	0	9 8	3	2000	2000	2000	0	0	0		000	0	000	0	2000	2000	1000	0		0	0	0	٤
똢	•	9	8/7	2778	3889	1667	833	0	256	0		0	1053	263	276	0	263	6239	276	789	0	;	250	250	0577	500	2250	1500	90	0	0	•	0 0	300	250	1250	4250	80	0	250		0	0	750	0
E	•	٥ :	44	1342	5299	1866	895	75	75	0		231	69.	382	169	231	308	8089	769	231	0	•	٥	285	/C81	1786	2357	1357	1857	0	0		0 6	3824	926	515	3676	368	0	0		0	22	362	,,,,,
5	į	Š,	0	2222	3889	3333	0	0	0	0		226	0	0	Ξ	0	0	8777	556	0	0	•	0	2857	/14	2143	2857	714	714	0	0		0 %	3889	Ξ	556	3333	556	0	0		0	0	226	
WA	•	Э (0	1991	833	7500	0	0	0	0		0	0	0	0	0	833	7500	1991	0	0	•	0	8	2000	000	3000	3000	0	0	0		0 6	0	90	1000	7000	0	0	0	•	0	0	1250	•
Υ		0	789	1316	3421	3684	789	0	0	0		734	0	0	0	0	588	8529	288	0	0		0	278	7777	1991	226	3333	1991	0	278		0 74	3235	1471	0	3529	0	0	0	•	0	0	263	1040
H		0	294	2059	4412	2647	288	0	0	0		0	625	0	625	0	0	5938	2500	312	0		0	0	3235	1764	2647	1764	288	0	0		0 %	3824	1764	0	3824	0	0	0	•	0	0	1250	3130
δ		0	417	1250	3333	2083	2917	0	0	0		0	417	0	0	0	1667	2999	1250	0	0		0	0	2083	833	417	4167	2500	0	0		0 6	3333	2083	0	3333	0	0		•	417	0	1667	5100
Ϋ́		0	333	2333	7997	2333	900	1000	333	0		0	714	357	0	0	0	6429	1786	714	0		0	8	8	1817	4545	0	1817	0	0		357	1429	0	1786	3571	2857	C		•	0	333	900	,,,,,,
Ħ		0	0	000	4000	4000	0	0	0	1000		0	800	0	2000	0	0	3000	2000	1000	0		0	0	0	3000	2000	0	2000	0	0		8 9	2000	2000	0	2000	0	0		•	0	0	0	200
ŊĠ		0	714	3213	2857	1786	1071	0	0	357		0	0	0	1154	1154	0	4615	2692	0	385		0	0	1071	2143	3571	1429	1786	0	0		0 0	92	92	2308	5385	92			•	0	0	357	;
TS		0	327	3213	4286	714	1071	0	0	357										0			0	714									0 25									0	0	0	9
SO	:	763	0	1579	4211	3158	276	0	0	263		0	333	333	1991	333	0	2000	2000	333	0		0	276	1316	2895	2105	1053	1842	263	0		0 %	1316	1053	1316	4737	789	-		>	0	0	276	1670
Allele		_	7	m	4	S	9	7	œ	6		-	7	60					00				_	7		4				∞	0		۰ -		4	v	۰	7	· 00	•	•	-	7	٣	•
	D1S407										D1S399											D2S273										D3S1537									D3S1545	2			

														ı			8	Ę	Ş		1	1	1		ı	ı	8	9	
Locus	Allele	8	TS		B		5	IJ	Ϋ́					9	LOCUS	VIICIC.	3	2 8			1	1	1	1	ı	П	١	اإ	
	12	1	0	1	1000	1	٥	888	526	1	ı	ı		18		- د	2 6	§ 5	0 7		•						90	0	
	13	_	714		0		0		526					0		۰ ۳	2895	2692	3571		• •			٠.			2750	2000	
	17	1316	1071	357	1000		833	0	0	0	0	362	263	0		4	3684	2000	3213	8	3929 2	86 2167	3824 520	5263 5000	0 3333	4143	5750	4000	
	15	_	1429		2000		417		0					0		ď	2368	385	1786							_	8	2000	
	16		357		0		0		0					0		9	789	0	357								0	2000	
	17		1429		0		0		0					0		7	0	0	357								0	0	
	18		0		0		0		276					0	D8S499														
	19		714		0		417		1842					0		-	0	385									0		
	20		0		0		0		0					0		7	0	0									0		
	21	_	357		0		5417	•	5789	_	٠.,			0			278	0									0		
	77		0		0		0		0					0		4	278	382									0		
D6S400																S	833	1154	1071	0	357	0	588 20	263 2500	0 714	1214	750	0	
	1	0	0		0		0	0	312					0		9	1389	92									9		
	2	0	0		2000		417	0	0					8		7	1389	1923									750		
	3	938	385		0		0	0	0				-	8		00	833	692									0		
	4	938	769		1000		833	294	0					0		6	833	385									0		
	5	938	1538		2000		0	0	0					8		9	1667	769									1750		
	9	938	385		0		0	0	312					0		=	1667	154									1250		
	7	2500	2308		0		833	588	312		_			0		12	833	2308			• • •		•				2000		
	∞	625	385		1000		0	294	0					8		13	0	0			•		•••				2500		
	6	312	769		0		0	0	0					8		14	0	0							0 0		0		
	2	625	385		0		1667	288	312	0	2778 1	324	789	8	D8S384														
	=	2188	3077		4000		6250	8235	8750	-	v	•		8		-	0	0									Š	0	
	17	0	0		0		0	0	0					0		7	263	0									3200	0	
1768303	!	1														3	2368	4286									3250	0	
1003393	-	263			c	0	0	312	263					0		4	3684	4643	-		-						2250	4000	
	,	236			0	357	0	0	263					8		2	2895	357									220	2000	
		526			2000	714	2500	2500	3158					0		9	276	0	357	0	1429	0 18	1875 294	2000	0 2500	429	250	0	
	4	9			1000	2143	3750	1250	526					8		7	0	714									0	1000	
		2105			3000	714	833	1562	1053	2000			_	00		∞	263	0									0	0	
	, v	5789			4000	2000	833	938	1842					0	D9S249												,	0	
	7	789			0	357	0	0	0		0	71	0	0		۰ ,	5789	7143									8 8	8 6	
	00	0	0	0	0	714	2083	3438	2895	1000				00		4 m	0	357	357	} •	· } •	0	0 0	0	0	0	0	0	
D7S620	-	•			9	c		c	c					0	D9S762	•	•	•									•	c	
	7	0	0	•	0	0	0	0	0	0	0	0	0	8		- 7	0	0	0	0	327	417 3	333 0	0	0 556	0	0	0	
		0	0		0	0	0	0	0					80		9	0	1154									0	0	
	4	1842	357		1000	1000	417	887	1842					00		4	1429	3077									2222	3000	
	•	2895	3571		1000	3667	4167	2000	4211		-			8		ς.	3571	2308	_		•						3889	2000	
	٠ ٧	3421	2857	-	3000	1667	3750	3235	3684					8		9	3571	1923									2778	3000	
	7	236	357		2000	1667	1991	288	263					8		7	714	1538			• • •	•	-				Ξ	2000	
	· 00	526	357		2000	1000	0	294	0					0		•	714	0									0	0	
	6	789	1786		0	1000	0	0	0					0	D10S526														
	10	0	714		0	0	0	0	0					0		-	563	0	0	0	0	0	625	0	0	227	263	0	
D7S623																													

æ	0	00	0009	000	0	0	0	0		0	0	0	0	٥	000	0 (-	, c		0	00	0	000	0		00	000	0	3000	2000	00	000	o (-	.	>	c	• •	90	2000	2	202
			3000														96 6											0											28			
				-				0									200										•	0											2029			
								0									35											Ξ									c			4286		
			833											•			3333										•	0	•								c	•	, 000			
			2940							0	0	263	368	842	632	68 5 68 5	1053	9 -	• •	0	0	0	263	0		312 875	875	0	375	0	250	0	312	0 (۰ د	9	278	623	2200	889	3 8	3
			1562 2	• •									• •		• •		1875												-										3824 2			
			2000	•													2083																						3750			
			• • •					0						• •			333																						. 692			
			3000														90											000											000			
								0		0	357	714	2143	321	714	1001	0 0	, [201	. 0	357	0	357	2143	714		357	1786		2857	0	1786	0	1429	357	1429	0	357		2143	3571		[0]
IS	357	2143	3929	2500	357	357	357	0		0	0	385	169	2308	169	<u> </u>	0 %	51	92	0	0	169	1154	0		00	2500	714	3929	0	2857	0	0	0 (o (0	•	•	2500	2000	3 5	357
ဒ္က	263	789	4737	2632	789	0	263	263									0 %									278	3 =	0	3889	0	3055	0	278	0	Š,	0	•	201	1316	3684		202
Allele	7	3	4	2	9	7	00	6		-	7	3	4	S	9	,	∞ 0	2 ۱	=	12	13	7	15	16		- ~	۰ ۳	4	8	9	7	∞	6	으 :	= :	12	-	٠,	4 ~	4	٠,	•
Locus									D14S119																D15S195											!	D16S485					
MB	0	2000	1000	0	0	0	0	0	0	0	1000	2000	2000	1000	0	1000	0	•	3750	2500	0	0	0	0	0	1250	1250	1250	> <	>	0	0	4000	1000	900	1000	0	0000	900	9	7007	
Ŧ	1842	2105	276	263	526	0	0	0	263	0	276	1579	526	1316	263	0	0	•	250	8	900	0	0	750	0	2000	3750	2000	2	>	0	0	Ξ	3889	3022	1944	0		3500	3750	06/7	
NE.	687	687	1364	8	1288	379	152	0	0	0	727	530	758	1591	1212	0	0	:	36.5	652	145	17	202	202	652	2029	7464	1594	6	>	0	92	833	409	2955	2045	0	;	9124	2000	107	
7	Ξ	0	226	226	Ξ	0	0	0	0	0	1667	0	0	3889	0	0	0	ì	55 56	1667	0	0	556	0	Ξ	Ξ	2222	ΞΞ	: `	>	0	0	2778	200	Ξ	Ξ	0	0	8889	25.	9	
Ψ	1000	0	0	0	0	0	1000	0	0	0	0	0009	900	0	1000	0	0	(0	0	1000	0	0	0	0	3000	8	2000	> <	>	0	90	300	300	2000	90	0	0	3 9	9	3	
٩	226	0	1053	0	263	789	0	0	0	0	226	2105	1579	2368	789	0	0	Č	278	1389	Ξ	0	0	0	278	1944	194	1667	6	>	0	0	1250	200	2500	1250	0	į	è °	223	6667	
5	625	1250	2188	312	0	312	0	0	0	0	938	1562	938	1250	0	0	0	Ş	88 88	794	288	0	0	0	294	1764	4412	9/11	2	167	0	0	0	2992	2333	9	900	;	<u>4</u>	3050	4007	
5	833	417	417	0	0	0	417	0	417	0	833	1667	1250	3333	417	0	0	•	455	2273	0	0	0	455	0	1817	2000	0 0	> <	>	0	0	2917	4583	2083	417	0		<u> </u>	8	ŝ	
ĕ	0	999	1333	333	0	0	333	0	333	999	99	999	1991	2333	90	0	0	į	ر د 0	357	714	714	0	0	357	1429	2143	2857	3 5	<u> </u>	٥.	382	1154	200	2308	1154	0	9	2000	2272	5/77	
R	0	0	1000	900	0	0	0	0	0	0	2000	90	90	2000	90	900	0	•	700 C	90	1000	0	0	0	0	3000	8	8 8	3 9	>	0	0	300	200	8	8	0	9	8	\$	3	
Š	0	0	0	1071	357	0	0	0	0	0	1786	1786	1071	1429	2143	357	0	•	714	357	0	0	0	357	357	2143	327	102		>	357	357	321	2857	6 43	1429	0	;	/143	2067	697	
2	0	769	385	385	382	0	0	385	0	0	1538	3077	1154	769	0	769	382	ģ	ğ 8	1154	385	0	0	0	0	1538	3077	1538	1	>	714	0	0	3571	2200	2857	327	9	853	3463	3407	
3	263	263	263	263	0	0	0	0	0	263	526	3158	1316	2105	0	1053	263	Ş	8 8 8	526	0	789	0	0	0	2105	1053	2895	747	3	263	0	1579	3158	2895	2105	0		7500	200	<u>x</u>	
Allele	7	e	4	2	9	7	∞	6	01	=	12	13	14	15	16	17	18	-	- 7	e	4	S	9	7	∞	6	2	= 5	7 [2	-	7	m	4	S	9	7		٦,	٠, ١	•	
Locus																		D10S516												D10S525							i i	HKASI				VWFII

9	0	0	0	0	0		0	2000	2000	1000	1000	1000	0	2000	000	-	2002	2000	3000	3000	0	0	0	0	0																		
FR	1500	520	750	250	0									0			90																										
				786						• •	-			0																													
M	2222	Ξ		929	111									0			0		٠.	• •			•																				
1				2000			0	0	0	3333	1999	0	0	0	0		• •																										
Ąſ	1053	1316	1579	0	789		263	1053	1842	2105	1421	1053	263	0	0	•	28°	764	412	764	0	0	764	0	0																		
	_	_	_	625						•				0			0		-																								
ర	833	1991	833	417	1250		0	417	1991	2917	3333	1250	417	0	0		833																										
S.	999	0	0	0	0		0	357	714	3213	2857	1786	0	357	714	•	0	1500	000	3000	90	200	0	0	0																		
Ħ	٥	0	0	1000	0		0	0	1000	3000	2000	1000	0	0	3000	•	0	2000	900	2000	0	3000	2000	0	0																		
2	ŀ	0	0	0	0		0	1071	3213	1429	2857	714	0	0	714	•	714	1786	3213	3571	0	714	0	0	0																		
TS	357	0	714	0	0		0	714	2500	2500	1786	1429	0	0	1071	•	357	2857	2500	2500	714	357	357	0	357																		
So	526	0	0	0	0		263	276	2632	2632	2105	1053	0	263	276	8	8 8	1579	4211	2632	526	263	0	263	0																		
Allele	∞	6	2	=	12		-	7	e	4	\$	9	7	00	6	•	- 7	(1)	. 4	5	9	7	00	6	2																		
Locus						D20S161										D20S428																											
9	3000	0		•	0	0	0	1000	7000	2000	0	0	0	0	0	c	0	2000	3000	1000	4000	0	0		0	1000	3000	000	1000	2000	0	0	000	000	0	0	c	3000	2000	2000	1000	2000	0
Æ	c	0		,	0	220	1500	200	3200	900	1750	0	0	0	0	-	278	1944	2000	1389	278	556	556		0	250	1250	750	750	0	0	0	9200	200	0	0	-	0	250	2250	000	1750	2000
里	c	145	72	!	217	0	1087	1522	3913	2101	1014	72	0	72	0	-	214	1143	6071	1571	286	429	786		286	714	357	53	200	0	0	0	6369	571	7	0	-	143	785	1714	1786	929	2357
1	ı							_	٠,	•				0			•	٠.	٠.														-			256							
×	1000	0	• •	•	0	0	0	3750	6250	0	0	0	0	0	0	•	•	0	10000	0	0	0	0		0	1667	833	833	0	833	0	0	2000	0	0	833	-	0	0	0	0	900	2000
ļ≤	c		· c	•	0	833	833	833	4167	3055	278	0	0	0	0	250	90	3158	5526	1053	0	0	0		0	0	263	0	0	0	0	276	2711	0	0	0	<	226	0	526	1579	1053	1579
핑	٦		• •	>	0	0	99	2000	6333	999	333	0	0	0	0	•	88	3235	5588	294	0	0	0		0	312	0	0	0	0	0	338	8750	0	0	0	•	312	0	312	625	1250	2188
5	٥	, c	• •	>	0	0	1364	1817	5455	1364	0	0	0	0	0	•	833	2500	6250	417	0	0	0		0	417	417	0	0	417	0	417	8333	0	0	0	•	417	833	1250	0	1667	833
8	٦	• •	•	>	0	0	333	000	2667	1000	3667	99	0	0	999	•	136	2273	3182	1817	88	455	0		0	0	80	200	0	200	900	200	2200	100	8	0	;	6	999	3333	8	2000	2333
Ä	١	•	•	>	0	0	0	2000	2000	000	0	000	0	000	0	•	0	0	000	1000	80	0	0		0	0	000	8	0	900	0	3000	2000	2000	0	0	•	2000	3000	2000	0	900	1000
1	1	> <									•			0		•	357	714	98/9	714	1429	0	0											٠.		0							101
1	П	•						-		-	٠.			357			0								0	357	357	1786	357	714	714	714	4286	714	0	0	•	o	3571	2143	107	1429	714
Ş	3 25	9	•	>				_		-	٠.			0		Š	\$ 88 8	288	5588	1471	1471	0	0		0	0	789	1579	789	0	1316	1316	2632	1579	0	0	Š	263	3684	2368	1053	263	1579
Allele	-	~ OI		^	-	7	٣	4	S	9	7	œ	6	2	=	•	- 7	· (*)	4	ν.	9	7	•		-	. 7	3	4	5	9	7	••	6	0	=	12	٠	- ~	۳ ۱	4	• •	9	7
I come				0178010	(100)											D18S930								D19S403													D19S400						

Figure AI Allele frequencies (×10,000) for 30 tetranucleotide repeat polymorphisms. SO = Sotho/Tswana; TS = Tsonga; NG = Nguni; BI = Biaka Pygmy; SA = San; C[‡] = Cambodian; CH = Chinese; JA = Japanese; MA = Malay; VI = Vietnamese; NE = northern European; FR = French; and MB = Mbuti Pygmy.

Locils	OS.	, E	N C	ä	A.S.	5	H	_ 4	¥ ¥	5	E	8	¥
178-350	\$25	-	101		٦	9223		7555	600	7857	4348	4737	
181-665	7637	7607	71/42		0000	6660	7041	0000	4167	7555	27.25	7750	
200-101	2601	777	107		6760	3333	3570	7500		1111	7084	2105	
107-01	203	417	1070	9	3143	5000	7647	2007		7143	6103	6500	000
180.400	0471	1071	6760	300	717	7060) o		1350	G (2766	600	
100-001	0//7	10/1	41/	7007	114	5	7.74	> ;	0071)	9/4	1001	> ;
178-648	4211	6154	3846	3000	1429	5833	5294	2000	0009	<i>L</i> 999	6118	6316	3000
251-590	929	1250	692	1000	385	1250	1176	1316	2000	625	3529	4062	1250
185-229	6316	4643	6071	8000	2000	1667	2000	5263	4167	2222	4191	3750	0006
171-808	1579	1786	2308	3000	2308	1667	1471	1667	4167	2222	3197	4474	2000
177-841	4737	4231	3929	7000	3462	2500	3125	4167	1667	3889	3712	3000	0006
177-108	3333	2000	4643	2000	3929	<i>L</i> 999	7059	7000	7500	7222	5448	2000	3000
187-676	4211	3077	2308	2000	1071	417	0	0	1667	1667	4071	3750	1000
285-016	10000	8846	10000	10000	9615	3333	4062	1944	2500	5556	5149	3333	10000
181-821	3158	3929	3571	4000	5357	5417	5625	2000	5833	3571	4130	3421	2000
180-411	7647	7307	8077	0009	7000	<i>L</i> 999	6176	6842	7000	7222	2959	7250	10000
250-584	1316	1071	1786	0	0	2500	294	938	0	0	143	0	0
178-649	4118	3462	3571	2000	4286	5417	5333	3750	4167	5556	6961	6229	3333
181-402	8333	8846	8571	0006	7857	2500	1471	1875	0	0	3551	5278	8750
182-559	2105	3077	2857	3000	1071	8750	7188	8947	0006	<i>1999</i>	6103	6229	7000
180-743	6259	5714	5357	0009	6071	3333	3235	2000	4167	3333	6071	2000	0006
196-856	3158	2143	1429	3000	1429	7500	5294	6176	7500	5556	4000	4750	0
177-381	9722	10000	9886	8750	2996	3750	2812	2632	7500	3889	3043	2000	10000
210-937	789	692	1429	0	0	3750	2059	2647	2000	1875	280	1500	3000
180-968	4722	2000	3462	2000	3571	3750	4062	4375	4000	1667	6562	9/19	2000
182-529	1053	1786	1154	3333	1071	3750	3235	2188	0	1250	2357	2632	0
192-312	5263	2000	2000	6250	6429	3750	3824	2222	833	4444	3623	5250	2000
196-905	2000	3462	2000	2000	0006	6250	5881	7500	5000	5625	4776	3684	0
181-184	4722	8169	3846	7000	6538	4583	3438	2940	3000	3750	2077	3421	2000
182-305	10000	10000	9643	10000	10000	9583	10000	9722	9167	10000	2186	9200	10000
185-174	8116	6538	<i>L</i> 999	2000	8182	5833	4375	4118	5000	2857	4609	5526	7500

Figure A2 Allele frequencies (×10,000) for 30 RSPs. Population abbreviations are given in the legend to fig. A1.

Population	Sequence (36 variable sites)	Population	Sequence (36 variable sites)
Sotho/Tennon	TAGATTIGACCARGCCAAGT	Nguni	TGAATTTGTTCTCGCACCC
Soulo/ I swalla	TASA A TETTASTICACIONA DE ATTASTA CONTRA RELATIONA DE LA CONTRA DE ATTASTA DE LA CONTRA DE ATTASTA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DE LA CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DE LA CONTRA DE LA CONTRA DE LA C	Nguni	TAGCTTTGTCCTCACGCTC
Sotho/18wana Sotho/Temana	TO A A TIMESTOTICE A CONTRACT A TOTAL A CONTRACT A CONTRACT A CONTRACT A TOTAL A TOTAL A CONTRACT A CONTRACT A TOTAL A CONTRACT A CONTRA	Nguni	TAGATTTGTCCCCGCGCTC
Sound/18wana	TYSA A THTHYST CHICAGO CONTOUR A THIGH THIS A CHICAGO CONTOUR A THIGH THIS A CHICAGO CONTOUR A THIGH THIS A THIRD A CHICAGO CONTOUR A THIRD A CHICAGO CONTOUR A CH	Nguni	TGAATTTTGCCCCCCGCACCC
Somo/ I swana	10AA11101C1C1OACC1C1A1OA11OA11ACAIA	Nguni	TGAATTTGCCCCCCGCACCC
Sotilo/15waiia	TAGGTTGTCTCAGGTTGTTTGAGTAGAGT	Nguni	TAGATITIGICCCCGCGCGCTC
Soulo/15walla	TGA A TITICOCOCOCOCOTA ATTACA	Biaka Pygmy	TAGATITGCCCTCGCACTC
Sotho/Tewana	TGAATTTGCCCCCCCCTATGATTACAGGT	Biaka Pygmy	TGGATTTGTCCCTGACCCC
Sotho/Tewana	TAGATTTGTCCCCCCCTACAATTGACTACAAGT	Biaka Pygmy	TGAATTTGTCTCCGCCCCC
Sotho/Tewana	TGAATTTGCCCCCCCCTTATGATTGATTACAGGT	Biaka Pygmy	TGAATTTGTCTCCGCCCCC
Sotho/Tewana	TGAATTTGCCCCCCCCCTGTGATTGATTACAGGT	Biaka Pygmy	TGAATTTGTCTCCGCCCCC
Sotho/Tswana	TAGCITTIGLCCTCACGCTCTATGATTGACTACAAGT	Mbuti Pygmy	TAAATTTGTCCTCGCACTC
Sotho/Tswana	TGAATTTGCCCCCGCACCCTATGATTGATTGCAAAT	Mbuti Pygmy	TGGATTTGTCCCCGCACTC
Sotho/Tswana	TGAATTTGCCCCCCCCCTTATGATTGATTACAAAT	Mbuti Pygmy	TGAATTTGTCCCCGCACTC
Sotho/Tswana	TGAATTTGCCCTCGCACCCCATGATTGATTACAAAT	Mbuti Pygmy	TGAATTTGCCCCTGCACCC
Sotho/Tswana	TGAATTTGCCCCCGCACCCTATGATTGATTACAAAC	Mbuti Pygmy	TGAATHGICICGACCC
Sotho/Tswana	TGAATTTGCCCCCGCACCCTATGATTGATTACGGGT	San	IGAALIIGIICCGCGCIC
Sotho/Tswana	TAGATTTGTCCCCGCGCTCTACAATTGACTACAAGT	San	IGAATITGCCCCGCACC
Sotho/Tswana	TAGCTTTGTCCTCACGCTCTATGATTGACTACAAGT	San	IGAATTIGCCCCGCGCCC
Sotho/Tswana	TGAATTTGTCTCTGACCCTTATGATTGATTACAAGT	San	IGAAIIIGCCCCGCGCCC
 Sotho/Tswana	TGAATTTGTCCCTGCACTCTATGATTGATTACAGGT	San	TGAATTIGCCCCCGCACCI
 Sotho/Tswana	TAGATITIGICCCCGCGCTCTACAATTGACTACAAGT	San	IGAATITGCCCCCGCACCI
Tsonga	TAGATITIGICCCCGCCTCTACAATTGACTACAAAT	San	IGAATIIGCCCCCGCGCCI
Tsonga	TGAATTTGCCCCCGCACTCTATGATTGATTACAAAT	San	TO A A TIME COCCOCCOCCOCCOCCOCCOCCOCCOCCOCCOCCOCCOC
Tsonga	TGAATTTGCCCCCCCCCTCTATGATTGATTACAAAT	San	IGAATTIGCCCCGCGCCC
Tsonga	TAGATITIGICCTCACGCTCTATGATTGACTACAAGT	San	IGAA I IGCCCCGCGCC
Tsonga	TGAATTTGTTCTCGCGCTCTATGATTGATTACAGGT	San	TGAATTTGCCCCGGGCGCCC
Tsonga	TGAATTTGCCCCCCCCACTCTATGATTGATTACAGGT	San	IGAAT HGCCCCGCACCI
Tsonga	TGAATTTGCCCCCCCCCTCTATGATTGATTACAGAT	San	IGAATTIGCCCCGCACC
Tsonga	TGAATITIGITICTCACGCTCTATGATTGATTACAGGT	San	TO A TITTA COCTOCO ACTOCO
Tsonga	TGAATTTGCCCCCGCACTCTATGATTGATTACAGGT	Cambodian	TO A ATTRICT CATOR ACTOR
Tsonga	TAGATTTGTCCCCGCGCTCTACAATTGACTACAAGT	Cambodian	TGA A TTTGCCCCCGCACTC
Tsonga	TGAATTTGCCCCCCCCCTTATGATTGATTACAAT	Cambodian	TGAATTTGTCCTCGCACTC
Tsonga	TGAATTTGTTCTCGCACCTTATGATTGATTACAGGT	Cambodian	TGAATTTGTCCTCGCACTC
Tsonga	TAGCITIGICCICACGCICIAIGAIIGACIACAGI	Cambodian	TGAATTTACCCTCGCACTC
Tsonga	TGAATTTGCCCCCGCACCCTATGATTGATTTACAGGT	Cambodian	TGAATTTGCCCTCGCACTC
Nguni	TGAATTTGCTCTCGCACCCTATGATTGATTACAAAT	Cambodian	TGAATTTGTCCTCGCACTC
Nguni	TGAATTTGCCCCCGCACCTTATGATTGATTACAAAT	Cambodian	TGAATTTATCCTCGCACCC
Nguni	TGAATTTIGCCCCCCCCCTATGATTGATTACAAAT	Cambodian	TGAATTTGTCCTCGCACTC
Nguni	TGAATTTGTTCTCCCCTCTATGATTGATTACAGGT	Cambodian	TGAATTTGTCCCCGCACTC
Nguni	TGAATTTGICCCGCACCCIATGATTGATTACAGT	Cambodian	TGAATTTGCCCCCCGCACCC
Nguni	TGAATTTGICTCTGAGCCTTATGATTGATTACAGG	Chinese	TAAATTTGCCCTCGCACTC
Nguni	TGAATTTGCCCCCGCACCCTATGATTGATTACAGG	Chinese	TAAATTTGTCCTCGCACTC
Nguni	TGAATTTACCCCTGCACCCTATGATTGATTACAGGT		

ini.	TGAATTTGTTCTCGCACCCTATGATTGATTACAGGT
ini.	TAGCTTTGTCCTCACGCTCTATGATTGACTACAAGT
ini	TAGATITIGICCCCCCCCTCTACAATIGACTACAAGT
ini	TGAATTTGCCCCCCCCCTATGATTGATTACAAAT
in.	TGAATTTGCCCCCCCCCTATGATTGATTACAGGT
in.	TAGATTTGTCCCCGCGCTCTATGATTGACTACAAGT
ka Pygmy	TAGATTTGCCCTCGCACTCCATGATTGACTACAAGT
ka Pygmy	TGGATTTGTCCCTGACCCCTATGATTGACTACAAGT
ka Pygmy	TGAATTTGTCTCCGCCCCTACGATTGATTACAAGT
ka Pygmy	TGAATTTGTCTCCGCCCCTACGATTGATTACAAGT
ka Pygmy	TGAATTTGTCTCCGCCCCTATGATTGATTATAGT
uti Pygmy	TAAATTTGTCCTCGCACTCTATGATTGACTACAAGT
uti Pygmy	TGGATTTGTCCCCCCCCTCTATGATTGATTACAGGT
uti Pygmy	TGAATTTGTCCCCGCACTCTACGATTGATTACAGGT
uti Pygmy	TGAATTTGCCCCTGCACCCTATGATTGATTACAGGT
uti Pygmy	TGAATTIGICICCGACCCCIACGATIGATTACAAGI
	TGAATTTGTTCCCGCGCTCTGTGATTGATTACAGGT
	TGAATTTGCCCCCCCCCTATGATTGATTACAGGT
	TGAATTTGCCCCCGCGCCTTATAATTGATTACAAAT
	TGAATTTGCCCCCCCCCCTTATGATTGATTACAAAT
	TGAATTTGCCCCCCGCACCTTATGATTGATTACAAAT
	TGAATTTGCCCCCCCCCTTATGATTGATTACAAAT
	TGAATTTGCCCCCCCCCCTTACAATTGATTACAAAT
	TGAATTTGCCCCCCCCCCTATGATTGATTACAAAT
	TGAATTTGCCCCCCCCCCTTATAATTGATTACAAAT
	TGAATTTGCCCCCCCCCCTTATAATTGATTACAAAT
	TGAATTTGCCCCCCCCCTATGATTGATTACAAAT
	TGAATTTGCCCCCCCCCTTATGATTGATTACAAAT
	TGAATTTGCCCCCCCCCTTATAATTGATTACAAAT
	TGAATTTGCCCCCCCCCCTTATGATTGATTACAAAT
nbodian	TGAATTTACCCTCGCACTCTATGATTGATTACAGGT
nbodian	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
nbodian	TGAATTTGCCCCCGCACCCTATGATTGATTACAGGT
nbodian	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
nbodian	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
nbodian	TGAATTTACCCTCGCACTCTATGATTGATTACAGGT
nbodian	TGAATTTGCCCTCGCACTCTATGATTGATTACAGGT
nbodian	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
nbodian	TGAATTTATCCTCGCACCCTATGATTGATTACAGGT
nbodian	TGAATTTGTCCTCGCACTCTATGGTTGATTACAGGT
nbodian	TGAATTTGTCCCCGCACTCTATGGTTGATTACAGGT
nbodian	TGAATTTGCCCCCGCACCCTATGATTGATTTACAGGT
inese	TAAATTTGCCCTCGCACTCTATGATTGATTACAGGT
inese	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT

Domilation	Carmence (36 unrinhla citae)	Pomilation	Comence (36 voriable cites)
r opulation	ocqueince (50 variable sites)	T Opumenou	adjustice (50 Valiable sites)
Chinese	TAAATITIGICCCCCCCICTATGATITACAGGT	Vietnamese	TGAATITIGCCCICGCACICIAIGATCGATTACAGGI
Chinese	TAAATTIGICICCGCACICIAIGATTGATTACAGGT	Vietnamese	IGAATITGICCICGCACICIAIGATIGATIACAGGI
Chinese	TAAATTTGTTCTCCCACTCTATGATTGATTACAGGT	Vietnamese	TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT
Chinese	TAAATTTGTCCCCCCCCTCTATGATTGATTACAGGT	Vietnamese	TGAATTTGTCCTCGCACTCTACGATTGATTACAGGT
Chinese	TGAATTTGTCCTCGCACTCCATGATTGATTACAGGT	Asian*	TGAATTTGTCCTCGCACTCTGTGATTGATTACAGGT
Chinese	TGAATTTGCTCTCGCACTCCATGATTGATTACAGGT	Asian*	TGAATTTGTCCTCGCGCTCTATGATTGATTACAGGT
Chinese	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT	Asian*	TGAATTTGTCTTCGCACTCTATGATTGATTACAGGT
Chinese	TGAATTTGTCCTCGCACTCCATGATTGATTACAGGT	Asian*	TGAATTTGCCCTCGCACTCTATGATTGATTACAGGT
Chinese	TGAATTTGTTCCCGCACTCCATGATTGATTACAGGT	Asian*	TGAATTTGTTCTCGCACTCCACAATTGATTACAGGT
Chinese	TGGATTTGTCCTCGCACTCTATGATTGATTACAGGT	Asian*	TGAATTTGTCCCCGCACTCCATGATTGATTACAGGT
Chinese	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT	Asian*	TGAATTTGTTCTCGCACTCTATGATTGATTACAGGT
Chinese	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT	Korean*	TGAATTTGTCCCCGCACTCTATGATTGGTTACAGGT
Chinese	TGAACCTGCCTTCGCACTCTATGATTGATTACAGGT	Korean*	TGAATTTGTTCTCGCACTCCATGATTGATTACAGGT
Chinese	TGAATTTGCCCTCGCACTCTATGATTGATTACAGGT	Asian*	TGAATTTGCCCTCGCACTCTATGATTGGTTACAGGT
Japanese	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT	Asian*	TGAATTTGTCCTCGCACTCCATGATTGATTACAGGT
Japanese	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT	Asian*	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGTCCTCGCACTCTACGATTGATTACAGGT	Asian*	TGAATTTGTCTTCGCACTCCATGATTGATTACAGGT
Japanese	TAAATTCGTCCCCGCACTCTATGATTGATTACAGGT	Asian*	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT	Asian*	TGAATTTGTCCTCGCGCTCTATGATTGATTACAGGT
Japanese	TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT	N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGTCCTCGCATTCTATGATTGATTACAGGT	N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT	N. European	TGAATTTGTTCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT	N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGCCCTCGCACTCCATGATTGATTACAGGT		TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGTCCCCGCACTCTATGATTACAGGT	N. European	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGTTCTCGCACTCTATGATTGATTACAGGT		TAAATTTGTCCTCGCACTCTATAATTGATTACAGGT
Japanese	TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT	N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TGAATTTGTCCCCGCACTCTATGATTACAGGT	N. European	TAAATTTGTCCTCGCACTCTATGATTAATTACAGGT
Japanese	TGAATTTGTTCTCGCACTCCATGATTACAGGT	N. European	TGAATTTGTCCTCACACTCTATGATTAATTACAGGT
Japanese	TGAATTTGCCCTCGCACTCTATGATTACAGGT	N. European	TGAATTTGCCCTCGCACTCTATGATTAATTACAGGT
Japanese	IGAATITIGICCICGCACICIAIGATIGATIACAGGT	N. European	TAAATTTGCCCTCGCACTCTATGATTGATTACAGGT
Japanese	1GAATTTGTCCTCGCACTCTATGATTTGATTTACAGGT		TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Japanese	TO A A TITUTO COLOCACIONAL DALI DALLA DALLA DE LA LA TITUTO DE LA		TAAATTTGTCCCCGCACTCTATGATTGATTACAGGT
Malay	100 A TITLE TO COLOR AND A TITLE TO A TITLE TO A TITLE TO A TITLE TO A TITLE A		TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Malay	TOWN THE TOTAL CONTROL OF THE ACTION OF THE		TAAATTTGTCCTCGAGCTCTATGATTTGATTACAGGT
Malay	10.4 A TITLE CONTROLL A LOCAL I DE A LITTLE A LA LITTLE CONTROLLA A TITLE CONTROLLA A LITTLE		TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT
Malay	IGAATTIGCCCTCGCACCTATGATTIGATTACAGGT	N. European	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Malay	1GAA111G11C1CGCAC1C1A1GA11GA111ACAGG1		TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Malay	10 AAIIIGICCCCCCACICIAIGACIGALIACAGGI	N. European	TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT
v iemamese	TO A A TITUTO COLLOCACIONAL TO A TITUTO		TAAATTTGTCCTCGCACCCTATGATTGATTACAGGT
Vietnamese	1GAA111GICCICGCACICIAIGA11GA11ACAGGI		TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Vietnamese	1GAA111GICCCGCACICIAIGA11GA11ACAGG1	N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
Vietnamese	10AA111GCCCCCGCACICCA1GA11GA11ACAGG1		TGAATTTGTCCTCGCGTCCTATAATTGATTACAGGT
		N. European	TAAATITIGICCTCGCACTCTATGATTGATTACAGGT

1 .	
	TAAATTTGTCCCCGCACTCTATGATTGATTACAGGT
N. European	TGAATTTGTCCTCACGCTCTACGATTGATTACAGGT
N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
N. European	TGAATTTGTCCTCGCACCCTATGATTGATTACAGGT
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	TAAATTTGTCCCCGCACTCCATAATTGATTACAGGT
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	IGAATITUTCCICGCUTCCIACAATIGATIACAGGI
N. European	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
N. European	164A1110CCC1CCCACICIA1OA110A1CACAG1
	TAAATTTGTTCCCCACTCTATGATTGATTACAGGT
	TGAATTTGTCCTCGCACCCTATGATTGATTACAGGT
	TAAATTTGTCCTCGCACTCCACGATTGATTACAGGT
N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
N. European	TAAATTTGTCCTCGCACCCTATGATTGATTACAGGT
N. European	TGAATTTGTCCTCGCACTCTATGGTTGATTACAGGT
	TAAATITIGICCTCGCACTCTAIGATIGATTACAGGT
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	TAAATTTGCCCTCGCACTCTATGATTGATTATAGGT
	TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT
	TGAATTTGTCCTCACACTCTATGATTGATTACAGGT
	TGAATTTGTTCCCGCACTCTATGATTGATTACAGGT
	IGAATITIGICCICACACICIAIGATIGATIACAGGI
	TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT
N. European	TAAATTIGICCICGCACICIAIGATIGATTACAGGT
N. European	TYPA A TITTICATION TO A
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT
N. European	TAAATITIGICCTCGCACTCTATGATTGATTACAGGT
	TAAATTTGTCCTCGCACCCTATGATTGATTACAGGT
	TAAATTTGTCCTCGCACCCTATGATTGATTACAGGT
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	TGAATTTGTCCCCGCACTCTATGATTGATTACAGGT
	TGAATTTGTTCTCGCACTCTATGATTGATTACAGGT
	TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT
	TGAATTTGCCCCCGCACTCTATGATTGATTACAGGT
N. European	TGAATTTGTCCTCGCACCCTATGATTGATTACAGGT
N. European	TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT

IGAATTTGCCCCCGCACCCTATGATTGATTACAGGT IGAATTTTGTCCCCGCACCCTATGATTGATTACAGGT

CAAATTTGTCCTCGCACTCTATGATTGATTACAGGT TGAATTTTGTCCTCGCACTCTATGATTGATTACAGGT

French French

French

French

TAAATTTGTCCTCGCACCCTATAATTGATTACAGGT TAAATITIGICCTCGCACTCTATGATTGATTACAGGT TGAATTTGTCCCCGCACTCTATGATTTGATTACAGGT TGAATTTGTTCTCGCACTCTATGATTGATTACAGGT TGAATTTGTTCTCGCACTCTATGATTGATTACAGGT TAAATTTIGTCCTCGCACTCTATGATTGATTACAGGT TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT TGAATTTGTCCTCGCACTCTATGATTGATTACAGGT TAGATITIGICCTCGCACTCTATGATTGATTACAGGT TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT TAAATTTGTCTCCGCACTCTATGATTGATTACAGGT TAAATTTGTCCTCGCACTCTATGATTGATTACAGGT

IGAATTTGTCCTCGCACTCTATGATTGATTACAGGT

Population N. European

French French

French French

French

French
French
French
French
French
French
French
French
French
French
French
French
French

Sequence (36 variable sites)

Figure A3 Mitochondrial D-loop sequence data for each individual in the study	a for each individual in the study
population. Only the 36 variable nucleotides are shown. The other nucleotides correspond	The other nucleotides correspond
to the Cambridge Reference Sequence (CRS). The variable sites are positions 72, 73, 93,	ble sites are positions 72, 73, 93,
95, 125, 129, 139, 143, 146, 150, 151, 152, 182, 185, 186, 189, 194, 195, 198, 199,	5, 186, 189, 194, 195, 198, 199,
200, 204, 207, 210, 212, 217, 228, 235, 236, 239, 241, 242, 244, 247, 263, and 265	11, 242, 244, 247, 263, and 265
of the CRS. Subjects marked with an asterisk (*) were used in the 3-population analyses	used in the 3-population analyses
but not in the 13-population analyses.	

References

- Anderson S, Bankier AT, Barrell BG (1981) Sequence and organization of the human mitochondrial genome. Nature 290:457-465
- Bamshad M, Fraley AE, Crawford MH, Cann RL, Busi BR, Naidu JM, Jorde LB. MtDNA variation in caste populations of Andhra Pradesh, India. Hum Biol (in press)
- Batzer MA, Stoneking M, Alegria-Hartman M, Bazan H, Kass DH, Shaikh TH, Novick GE, et al (1994) African origin of human-specific polymorphic *Alu* insertions. Proc Natl Acad Sci USA 91:12288–12292
- Bell GI, Karem JH, Rutter JR (1981) Polymorphic DNA region adjacent to the 5' end of the human insulin gene. Proc Natl Acad Sci USA 78:5759-5763
- Birky CW, Maruyama T, Fuerst P (1983) An approach to population and evolutionary genetic theory for genes in mitochondria and chloroplasts, and some results. Genetics 103:513-527
- Bowcock A, Cavalli-Sforza LL (1991) The study of variation in the human genome. Genomics 11:491–498.
- Bowcock AM, Kidd JR, Mountain JL, Hebert JM, Carotenuto L, Kidd KK, Cavalli-Sforza LL (1991) Drift, admixture, and selection in human evolution: a study with DNA polymorphisms. Proc Natl Acad Sci USA 88:839–843
- Bowcock AM, Ruiz-Linares A, Tomfohrde J, Minch E, Kidd JR, Cavalli-Sforza LL (1994) High resolution of human evolutionary trees with polymorphic microsatellites. Nature 368:455-457
- Cann RL, Stoneking M, Wilson AC (1987) Mitochondrial DNA and human evolution. Nature 325:31-36
- Cavalli-Sforza LL, Menozzi P, Piazza A (1994) The history and geography of human genes. Princeton University Press, Princeton.
- Cavalli-Sforza LL, Piazza A, Menozzi P, Mountain J (1988) Reconstruction of human evolution: bringing together genetic, archaeological, and linguistic data. Proc Natl Acad Sci USA 85:6002-6006
- Deka R, Chakraborty R, Ferrell RE (1991) A population genetic study of six VNTR loci in three ethnically defined populations. Genomics 11:83-92
- Deka R, Jin L, Shriver MD, Yu LM, DeCroo S, Hundrieser J, Bunker CH, et al (1995) Population genetics of dinucleotide (dC-dA)_{n•}(dG-dT)_n polymorphisms in world populations. Am J Hum Genet 56:461–474
- Di Rienzo A, Peterson AC, Garza JC, Valdes AM, Slatkin M, Freimer NB (1994) Mutational processes of simple-sequence repeat loci in human populations. Proc Natl Acad Sci USA 91:3166-3170
- Edwards A, Hammond HA, Jin L, Caskey CT, Chakraborty R (1992) Genetic variation at five trimeric and tetrameric tandem repeat loci in four human population groups. Genomics 12:241-253
- Excoffier L (1990) Evolution of human mitochondrial DNA: evidence for departure from a pure neutral model of populations at equilibrium. J Mol Evol 30:125–139
- Felsenstein J (1993) PHYLIP (Phylogeny Inference Package) version 3.5c. Department of Genetics, University of Washington, Seattle
- Harpending H, Relethford J, Sherry S. Methods and models for understanding human diversity. In: Boyce AJ, Mascie-

- Taylor CGN (eds) Molecular Biology and Human Diversity. Cambridge University Press, London (in press)
- Horai S, Hayasaka K, Kondo R, Tsugane K, Takahata N (1995) Recent African origin of modern humans revealed by complete sequences of hominoid mitochondrial DNAs. Proc Natl Acad Sci USA 92:532-536
- Horai S, Kondo R, Nakagawa-Hattori Y, Hayashi W, Sonoda S, Tajima K (1993) Peopling of the Americas, founded by four major lineages of mitochondrial DNA. Mol Biol Evol 10:23-47
- Jin L, Chakraborty R. Population structure, stepwise mutations, heterozygote deficiency and their implications in DNA forensics. Heredity (in press)
- Jorde LB (1980) The genetic structure of subdivided human populations: a review. In: Mielke JH, Crawford MH (eds) Current developments in anthropological genetics, Vol. 1. Plenum, New York, pp 135–208
- Kidd JR, Black FL, Weiss KM, Balazs I, Kidd KK (1991) Studies of three Amerindian populations using nuclear DNA polymorphisms. Hum Biol 63:775-794
- Kimura M (1980) A simple model for estimating evolutionary rates of base substitutions through comparative studies of nucleotide sequences. J Mol Evol 16:111-120
- Mantel N (1967) The detection of disease clustering and a generalized regression approach. Cancer Res 27:209-220
- Merriwether DA, Clark AG, Ballinger SW, Schurr TG, Soodyall H, Jenkins T, Sherry ST, et al (1991) The structure of human mitochondrial DNA variation. J Mol Evol 33:543–555
- Mountain JL, Cavalli-Sforza LL (1994) Inference of human evolution through cladistic analysis of nuclear DNA restriction polymorphisms. Proc Natl Acad Sci USA 91:6515–6519
- Nei M (1987) Molecular evolutionary genetics. Columbia University Press, New York
- Nei M, Livshits G (1989) Genetic relationships of Europeans, Asians and Africans and the origin of modern Homo sapiens. Hum Hered 39:276–281
- Nei M, Livshits G, Ota T (1993) Genetic variation and evolution of human populations. In: Sing CF, Hanis CL (eds) Genetics of cellular, individual, family, and population variability. Oxford University Press, New York, pp 239–252
- Nei M, Roychoudhury AK (1982) Genetic relationship and evolution of human races. Evol Biol 14:1-59
- (1993) Evolutionary relationships of human populations on a global scale. Mol Biol Evol 10:927-943
- O'Brien E, Rogers AR, Beesley J, Jorde LB (1994) Genetic structure of the Utah Mormons: a comparison of results based on DNA, blood groups, migration matrices, isonymy, and pedigrees. Hum Biol 66:743-759
- Relethford JH, Harpending HC (1994) Craniometric variation, genetic theory, and modern human origins. Am J Phys Anthropol 95:249–270
- Rogers AR. Genetic evidence for a Pleistocene population explosion. Evolution (in press)
- Rogers AR, Harpending HC (1992) Population growth makes waves in the distribution of pairwise genetic differences. Mol Biol Evol 9:552-569
- Rogers AR, Jorde LB (1995) Genetic evidence on the origin of modern humans. Hum Biol 67:1-36
- Saitou N, Nei M (1987) The neighbor-joining method: a new

- method for reconstructing phylogenetic trees. Mol Biol Evol 4:406-425
- Sherry ST, Rogers AR, Harpending H, Soodyall H, Jenkins T, Stoneking M (1994) Mismatch distributions of mtDNA reveal recent human population expansions. Hum Biol 66:761-775
- Shriver MD, Jin L, Boerwinkle E, Chakraborty R (1995) A novel measure of genetic distance for highly polymorphic tandem repeat loci. Mol Biol Evol 12 (in press)
- Shriver MD, Jin L, Chakraborty R, Boerwinkle E (1993) VNTR allele frequency distributions under the stepwise mutation model: a computer simulation approach. Genetics 134:983–993
- Smouse PE, Long JC, Sokal RR (1986) Multiple regression and correlation extensions of the Mantel test of matrix correspondence. Syst Zool 35:627-632
- Spuhler JN (1993) Population genetics and evolution in the genus *Homo* in the last two million years. In: Sing CF, Hanis CL (eds) Genetics of cellular, individual, family, and population variability. Oxford University Press, New York, pp 262–297
- Spurdle AB, Jenkins T (1992) The Y chromosome as a tool for studying human evolution. Curr Opin Genet Dev 2:487–491 Stoneking M (1993) DNA and recent human evolution. Evol Anthropol 2:60–73
- Stoneking M, Jorde LB, Bhatia K, Wilson AC (1990) Geo-

- graphic variation of human mitochondrial DNA from Papua New Guinea. Genetics 124:717–733
- Sykes BC (1983) DNA in heritable disease. Lancet 2:787–788 Tajima F (1989) Statistical method for testing the neutral mutation hypothesis by DNA polymorphism. Genetics 123:585–595
- Takahata N, Palumbi SR (1985) Extranuclear differentiation and gene flow in the finite island model. Genetics 109:441–457
- Templeton AR (1993) The "Eve" hypothesis: a genetic critique and reanalysis. Am Anthropol 95:51-72
- Vigilant L, Pennington R, Harpending H, Kocher TD, Wilson AC (1989) Mitochondrial DNA sequences in single hairs from a southern African population. Proc Natl Acad Sci USA 86:9350-9354
- Vigilant L, Stoneking M, Harpending H, Hawkes K, Wilson AC (1991) African populations and the evolution of human mitochondrial DNA. Science 253:1503-1507
- Watkins WS, Bamshad MJ, Jorde LB. Population genetics of trinucleotide repeat polymorphisms. Hum Mol Genet (in press)
- Weber JL, Wong C (1993) Mutation of human short tandem repeats. Hum Mol Genet 2:1123-1128
- Wright S (1965) The interpretation of population structure by F-statistics with special regard to systems of mating. Evolution 19:395-420